

The Spot 42 RNA: A regulatory small RNA with roles in the central metabolism

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The Spot 42 RNA is a 109 nucleotide long (in *Escherichia coli*) noncoding small regulatory RNA (sRNA) encoded by the *spf* (spot forty-two) gene. *spf* is found in gamma-proteobacteria and the majority of experimental work on Spot 42 RNA has been performed using *E. coli*, and recently *Aliivibrio salmonicida*. In the cell Spot 42 RNA plays essential roles as a regulator in carbohydrate metabolism and uptake, and its expression is activated by glucose, and inhibited by the cAMP-CRP complex. Here we summarize the current knowledge on Spot 42, and present the natural distribution of *spf*, show family-specific secondary structural features of Spot 42, and link highly conserved structural regions to mRNA target binding.

Introduction

The *spf* gene is highly conserved in *Escherichia*, *Shigella*, *Klebsiella*, *Salmonella* and *Yersinia* (genera) within the *Enterobacteriaceae* family.¹ In *E. coli* the *spf* gene is flanked by *polA* (upstream) and *yihA* (downstream),^{2,3} and a CRP binding sequence and -10 and -35 promoter sequences are found upstream of *spf*. *spf* is also highly conserved within the *Vibrionaceae* family, and was recently identified in 76 *Vibrionaceae* genomes that were available at that time (e.g., *Vibrio*, *Aliivibrio*, *Photobacterium* and *Grimontia* genera).⁴ In e.g., *Vibrio cholerae*, *Vibrio vulnificus*, *Aliivibrio fischeri* and *A. salmonicida* the *spf* gene is flanked by *polA* (upstream) and a sRNA gene encoding the novel VSsrna24 RNA (downstream).

Spot 42 was first described in 1973 as an unstable RNA species of 109 nucleotides in *E. coli*.^{5,6} It was discovered by polyacrylamide gel electrophoresis and 2-D fingerprinting in an attempt to study the accumulation of small RNAs in *E. coli* during amino acid starvation. In these experiments the electrophoretic mobility of Spot 42 was similar to that of 5S rRNA. In 1979, Spot 42 was reported to accumulate under growth in the presence of glucose (i.e., when adenosine 3',5'-cyclic monophosphate (cAMP) is low).^{7,8} During growth with a non-glucose carbon source (i.e., when cAMP concentrations are high) Spot 42 concentrations were significantly lower. Later experiments showed that overexpression of Spot 42 (tenfold increase) resulted in impaired growth and lowered ability to adapt to shifts to richer media.⁹ Further, shift from glucose to succinate as the carbon source resulted in a long lag period and slow growth rate, the reason for the abnormal responses was caused by an elevated number of excessive Spot 42 RNA gene products rather than excess of the gene itself. A deletion study of *spf* in *E. coli* cells resulted in viable *spf* null mutants, which indicated that Spot 42 was non-essential, at least under controlled lab conditions.¹⁰

It was for some years unclear if the function of Spot 42 was mediated through the 109 nucleotide RNA itself or if the function was mediated through the 14 amino acids long peptide which is hypothetically encoded from within the sRNA sequence. This confusion was based on the observation that Spot 42 contains structural features similar to other non-coding RNAs found in *E. coli* (such as 6S RNA and lambda bacteriophage), as well as features that are typically found in mRNAs (i.e., polypurine sequence followed by AUG, 14 amino acids codons and an UGA termination codon).⁷ Using a filter binding assay and other methods Rice et al. showed that Spot 42 is not an mRNA.¹¹ In this approach the affinity between Spot 42 and the 70S ribosome was tested. Here, Spot 42 showed very inefficient binding to purified 70S ribosomes, which lead to the conclusion that the function of Spot 42 is mediated by the RNA itself.

The direct responsiveness of Spot 42 levels to glucose and cAMP is due to repression of *spf* expression by a cAMP-CRP (cAMP-receptor protein) complex.² The reduction of Spot 42 in cells grown in secondary carbon sources is a result of binding of the cAMP-CRP complex to the *spf* promoter, which negatively regulates transcription of Spot 42. Later, the proximity of *spf* to *polA* (gene encoding DNA polymerase I) led Dahlberg and co-workers to test whether the products of these genes could influence each other.¹² They found that by reducing levels of Spot 42, either by deletion of *spf* or by manipulating the growth conditions, the DNA pol A activity was reduced. The underlying mechanism for this observation remains however unknown.

Spot 42 can interact directly with mRNA targets through base pairing. The first Spot 42 target was discovered by Møller et al., who showed that Spot 42 specifically binds to a short

63 complementary region at the translation initiation region of *galK* (encodes a galactokinase)
64 mediated through binding of the posttranscriptional regulator Hfq.¹ *galK* is the third gene in the
65 galactose operon, which contains four genes (*galETKM*) and produces a polycistronic mRNA.
66 Spot 42 mediates discoordinate expression of the *gal* operon (i.e., the individual genes in the
67 operon are not similarly expressed) by binding to the *galK* Shine-Dalgarno region, thereby
68 blocking ribosome binding and translation of the *galK* gene. The physiological significance of the
69 discoordinate expression is unclear, but suggests that Spot 42 plays a role in fine-tuning gene
70 expression to optimize the utilization of carbon sources. Recently, Wang et al. showed that Spot
71 42 represses expression of *galK* through direct binding to the 5' end of the *galK* mRNA, and also
72 mediates transcription termination of *galT* in the *galT-galK* junction.¹³

73 Beisel and Storz demonstrated with microarray analysis and reporter fusions that Spot
74 42 plays a broader role in metabolism by regulating at least 14 operons.¹⁴ These operons contain
75 a number of genes involved in uptake and catabolism of non-favored carbon sources. During
76 overexpression of Spot 42 sixteen different genes showed consistently twofold reduced or
77 elevated levels of mRNA. The identified reduced genes are mostly involved in central and
78 secondary metabolism, as well as uptake and catabolism of non-preferred carbon sources and
79 oxidation of NADH. In 2012 Beisel et al. performed computational target analysis using the
80 three conserved regions of Spot 42 as input. Compared to when using full-length Spot 42
81 sequence as input the target identification was improved and additional targets were revealed.¹⁵
82 The target analysis combined with assays of reporter fusions identified seven novel Spot 42
83 mRNA targets, all involved in catabolite repression. Mutational analysis showed that the
84 interactions of the three conserved regions of Spot 42 are critical in target regulation and that
85 regulation through multiple conserved regions of Spot 42 as well as increased base-pairing in
86 these regions strengthen the target regulation.

87 The evolution of sRNAs in *E. coli* and their regulatory interactions with mRNAs was
88 recently studied using computational methods.¹⁶ Compared to cis-acting sRNA and other non-
89 coding RNA (housekeeping RNA), trans-acting sRNA was the latest to appear in evolution.
90 Furthermore, after Enterobacteriales diverged into a separate lineage within gamma-
91 proteobacteria, the trans-acting sRNAs likely appeared in relatively high numbers compared to
92 the cis-acting sRNAs that evolved more evenly among all orders within gamma-proteobacteria.
93 The evolutionary age of 15 sRNAs and 49 corresponding sRNA-mRNA interactions were
94 examined. Here, Spot 42 was found to be the most ancient sRNA. Of the six Spot 42 mRNA
95 targets considered, only two (*xylF* and *galK*) evolved before Spot 42, albeit all the
96 complementary mRNA binding sites appeared after Spot 42.

The observation that *A. salmonicida* contains the *spf* gene (which encodes the Spot 42 RNA), but lacks the *galK* operon (the natural Spot 42 target in *E. coli*), have inspired scientists to study the role of Spot 42 in this fish pathogen.⁴ *A. salmonicida* is unable to utilize galactose (lacks *gal* operon) in minimal medium and addition of galactose has little effect on the growth rate. When cells are grown in glucose the level of Spot 42 is increased 16-40 fold, but is in contrast decreased threefold when cAMP is added, indicating that Spot 42 have similar roles as in *E. coli*, i.e., in carbohydrate metabolism. It has been hypothesized that Spot 42 works in concert with a novel sRNA gene, called *VSsrna24*, located 262 nt downstream of *spf*. The *VSsrna42* RNA is approximately 60 nt in length and has an expression pattern opposite to that of Spot 42. Furthermore, in a *spf* deletion mutant a gene encoding a pirin-like protein was upregulated 16 fold. Pirin has key roles in the central metabolism by regulating the activity of pyruvate dehydrogenase E1 and therefore select whether pyruvate will be fermented, or subjected to respiration through the TCA cycle and electron transport.

Although the Spot 42 RNA was discovered more than 40 years ago there are still a number of unanswered question related to this highly interesting RNA, e.g.: What is the natural distribution of the Spot 42 gene (*spf*) in Bacteria? What is the complete set of biological roles of Spot 42, and does Spot 42 play the proposed key role in the central metabolism? How does Spot 42 interact with its apparently many mRNA targets? In this work we have summarized the current literature on Spot 42, and extended this knowledge by surveying the known natural distribution of *spf*, we have identified family-specific structural features of Spot 42, and evaluated if highly conserved structural regions can be linked to mRNA binding.

Results

***spf* is restricted to 5 orders of gamma-proteobacteria**

The distribution of *spf* in nature is shown in **Fig. 1**. The basis for the figure was available nucleotide sequences of *spf* included in the Rfam database (677 sequences), and *spf* sequences identified in this study by using the Blastn server and *spf* sequences from selected taxa as queries. All previously known cases of *spf* originate from gamma-proteobacteria, and after fruitless searches in all other domains of Bacteria we therefore concentrated our efforts on specific searches within gamma-proteobacteria, both by using *spf* sequences from the closest neighbors, and by manual inspection of the known genic location of *spf*, i.e., in the intergenic region between *polA* and *engB*. The result of our search was finally mapped onto a phylogenetic tree generated using the iTOL web service.

The result show that *spf* is exclusively found in five orders of gamma-bacteria, i.e., in Enterobacteriales, Aeromonadales, Alteromonadales, Vibrionales and Chromatiales. These orders, except Chromatiales, share the same closest common ancestor (arrow in **Fig. 1**), and constitutes a clade. *spf* has still not been found in Pasteurellales, which is likely due to that Pasteurellales genomes are underrepresented in the European Nucleotide Archive (ENA) compared to e.g., the sister Enterobacteriales. We suspect that *spf* will be discovered in Pasteurellales as more genomes are being sequenced. In addition to known cases of *spf* our Blastn search revealed previously unreported cases within genera of Enterobacteriales and Alteromonadales. In Enterobacteriales *spf* was identified in the genera *Morganella* and *Raoultella*, as well as in draft genomes of *Budvicia*, *Cedecea*, *Hafnia*, *Leminorella*, *Plesimonas* and *Yokenella*. And, in genera where *spf* was already known to occur, *spf* was in this work identified in *Enterobacter radicincitans* and *Escherichia blattae*. Similarly, in Alteromonadales *spf* is found in the five families *Ferrimonadaceae*, *Shewanellaceae*, *Moritellaceae*, *Pseudoalteromonadaceae* and *Alteromonadaceae*, and *spf* was in this study identified in the three genera *Glaccola*, *Alteromonas* and *Pseudoalteromonas* by our blast searches, whereas *spf* was found in *Moritella viscosa* by manual inspection of the intergenic region *polA/engB*. Interestingly, in Chromatiales, *spf* is exclusively found in the genera *Rheinheimera* and *Arsukibacterium*, which is represented in ENA by six and two available draft genomes, all containing *spf*. Given that the phylogeny as shown in **Fig. 1** is correct then it is tempting to speculate that *spf* was acquired by lateral transfer, perhaps from a donor within the clade marked by an arrow in **Fig. 1**.

We also wanted to answer the following question: Is *spf* optional or ubiquitous within the individual orders and families? Spot 42 appears to play central roles in the carbohydrate metabolism, and we therefore hypothesized that it might be present in all representatives of the same order, family or genus once it has been identified in one genome. To answer this question we used the list of complete bacterial genomes found at the NCBI Genomes resource (<http://www.ncbi.nlm.nih.gov/genome/>), and searched for presence of *spf* in all representatives of the current orders, families and genera. Our result show that *spf* is found in 699 of 741 complete genomes distributed among 34 genera (a detailed list is provided in **Table S1**). *spf* is missing in representatives of the two genera *Glaccola* and *Pseudoalteromonas* of Alteromonadales. In both of these genera *spf* is found in one of three complete genomes. All three genomes of *Glaccola* have the same genic organization with *polA* and *engB* as neighbors (*spf* is usually located between these two genes). In *Pseudoalteromonas*, *spf* is only found in one genome, i.e., in *Pseudoalteromonas atlantica*, where *polA* and *engB* are located next to each other. The two other genomes with no *spf* have a different genic organization (synteny) at this region. Finally, *spf* has not been found in any of the complete genomes within the following genera:

Buchnera, *Candidatus* Moranella, *Candidatus* Riesia and *Wigglesworthia* (from Enterobacteriales), *Oceanomoas* and *Tolumonas* (from Aeromonadales), *Marinobacter*, *Sacchrophagus*, *Colwellia*, *Idiomarina* and *Psychromonas* (from Alteromonadales), and all genera of Chromatiales (i.e., *spf* found in 6 draft genomes of the genus *Rheinheimera* and 2 draft genomes of *Arsukibacterium*). In summary, of a total of 741 genomes from the 5 orders *Enterobacteriales*, *Aeromonadales*, *Alteromonadales*, *Chromatiales* and *Vibrionales*, 699 complete genomes contain *spf*, whereas 42 lack *spf*. The result is in agreement with conserved, but not necessarily indispensable roles of *spf*.

The Spot 42 RNA consensus secondary structure

We next mapped the level of identity among all known *spf* sequences (120 in total when redundant sequences have been removed) onto a consensus secondary structure model of Spot 42 (based on structure probing by Møller et al.¹) to find clues to possible structural regions that might be important for target identification and interaction, in general (**Fig. 2**). The Spot 42 RNA consists of one long hairpin structure located at the 5' end (from now on referred to as the 5' hairpin; 45–59 nt in length), and a second smaller hairpin separated from the 5' hairpin by a 9 - 20 nt long single-stranded region. In addition, a rho-independent terminator is located immediately downstream of the second hairpin. Structural regions of Spot 42 from the families *Vibrionaceae*, *Aeromonadaceae* and *Shewanellaceae* differ from the general “consensus” and are shown in separate boxes in **Fig. 2**. The sRNA gene is, in general, highly conserved with 76 of 108 positions (when using the “consensus” sequence as the reference) being 80–100% identical across all orders (shown as uppercase bold letters in **Fig. 2**). Notably, the 5' hairpin is highly conserved, i.e., 80–100% identity from positions 1–41, which indicate that these positions are interesting candidates for having general roles in target binding, perhaps with the terminal loop functioning as the seed sequence. The single-stranded region separating the 5' hairpin and the second hairpin is less conserved, with 80–100% identity in three positions and 60–79% identity in six positions, and is therefore perhaps less likely to have general roles in target recognition. *spf* is as expected most conserved within families. The *Shewanellaceae* *spf* differs most from the “consensus”. Here, the 5' hairpin contains two bulges with eight additional nt (inserted between pos. 39 and pos. 47). The *Vibrionaceae* and *Aeromonadaceae* sequences also differ to some extent from the “consensus”. In summary, Spot 42 is a highly conserved sRNA across five orders. The 5' hairpin represents the most conserved region and is therefore expected to have general roles in target recognition and interaction.

Spot 42 structure conservation and potential base pairing with targets

We next wanted to investigate if the highly conserved nucleotide positions of Spot 42 (as described above) are implicated in target binding (i.e., base-pairing between Spot 42 and mRNA target). Interactions between Spot 42 and *galK* mRNA has been determined using structure probing,¹ whereas potential base-pairing to other targets is based on bioinformatics predictions followed by experimental work.^{4,14,15}

Fig. 3 shows schematically potential base-pairing between Spot 42 and experimentally verified mRNA targets for the following genes: *galK*, *pirin*, *fucI*, *xylF*, *sthA*, *gltA*, *srlA*, *nanC*, *paaK*, *ascF*, *caiA*, *fucP*, *atoD*, *puuE* and *nanT*. Interestingly, for all except two genes (i.e., *sthA* and *fucP*) the most conserved region of the 5' hairpin (i.e., pos. 1-41) can potentially participate in extensive base-pairing with the corresponding mRNAs. This suggests that the 5' hairpin, is essential for target recognition and binding. Moreover, the first six positions of Spot 42 (5' single stranded region) can potentially base-pair with ten of fifteen targets (*galK*, *pirin*, *fucI*, *xylF*, *gltA*, *nanC*, *paaK*, *ascF*, *atoD* and *nanT*), and the terminal loop of the 5' hairpin can base-pair with eight of fifteen targets (*galK*, *pirin*, *fucI*, *xylF*, *srlA*, *caiA*, *puuE* and *nanT*). The second hairpin is only partly conserved. In agreement with this observation base-pairing with targets are rarer and only observed for two targets (*galK* and *pirin*). This is in agreement with results from Beisel et al.¹⁵ Using three unstructured regions (the 5' single stranded region, the 5' hairpin and the single-stranded region separating the hairpins) as input during computational target identification, they improved identification of direct targets, compared to when using the full-length sequence of Spot 42. In summary, highly conserved nucleotide positions of Spot 42 have the potential to participate in extensive base-pairing with known mRNA targets.

sRNA genes in the intergenic region downstream of *polA*

Interestingly, *spf* is not the only sRNA gene located in the intergenic region downstream of *polA* (see **Fig. 4**). In *Vibrionaceae* a gene encoding the sRNA VSsrna24 is located approximately 600 nt downstream of *spf*. Expression of VSsrna24 is repressed by glucose, and is hypothesized to have roles in the central carbohydrate metabolism.⁴ The sRNAs sX13,¹⁷ ErsA¹⁸ and Smr7C,^{19,20} are found in *Xanthomonadaceae*, *Pseudomonas* and Rhizobiales, respectively, but neither has the same function or structure as Spot 42. sX13 and Smr7C share secondary structure features comprising three stem-loops with C-rich motifs and are Hfq-independent.^{17,21} ErsA is Hfq-mediated and regulated by sigma factor 22, in contrast to Spot 42 that is dependent on sigma

factor 70. If any of these four sRNA genes originates from a common ancestral gene or not is currently unknown.

Concluding Remarks

We have conducted a survey on Spot 42 RNA in order to learn about its natural distribution, conservation patterns, and mRNA target recognition. We demonstrated that Spot 42, which was first identified in *E. coli* (Enterobacteriales), is also common in four other orders, i.e., Aeromonadales, Alteromonadales, Chromatiales and Vibrionales. Using blastn analysis we discovered novel *spf* sequences. Of a total of 741 complete genomes from the 5 orders Enterobacteriales, Aeromonadales, Alteromonadales, Chromatiales and Vibrionales, 699 genomes contain *spf*. Furthermore, a total of 30 draft genomes distributed among 11 genera (from all orders except Aeromonadales) contain *spf*. As shown in **Fig. 1**, within gamma-proteobacteria, Aeromonadales, Alteromonadales, Enterobacteriales and Vibrionales share the same last common ancestor, whereas Chromatiales does not, which suggest that *spf* was introduced into Chromatiales by lateral transfer by a donor from the clade marked by an arrow. We made a consensus secondary structure model of Spot 42 based on all known *spf* sequences and compared this to a schematically figure showing potential base-pairing between Spot 42 and known mRNA targets. Our results show that highly conserved nucleotide positions, in general, have potential to participate in extensive base-pairing with target mRNAs. This is in agreement with an earlier study by Beisel et al. which suggested that the strength of Spot 42 regulation is directly dependent on the number of nucleotides and the number of highly conserved structural regions which are involved in base-pairing between Spot 42 and its target.¹⁵

It is intriguing to us that although Spot 42 was discovered more than 40 years, there are still many unanswered questions. As more sequence data are being produced from high-throughput sequencing techniques and better tools and search algorithms are being developed, the known natural distribution of *spf* will certainly expand to new orders, families and genera (and perhaps phyla). And detailed knowledge on target recognition (other than *galk*) and roles in cellular processes will come from functional and bioinformatics studies. One particularly interesting aspect of Spot 42 is its apparent central role (via pirin) in the central metabolism by directing pyruvate towards fermentation or respiration through the tricarboxylic acid (TCA) cycle and electron transport.

Materials and Methods

Homology search

All previously known *spf* sequences were retrieved from Rfam (<http://rfam.sanger.ac.uk/family/RF00021>).²² Blastn searches in all domains of Bacteria were performed using *spf* sequences from 43 selected taxa as query sequences. All complete bacterial genomes found at the NCBI Genomes resource (<http://www.ncbi.nlm.nih.gov/genome/>) were checked for the presence of *spf*. More thorough blastn searches were performed in gamma-proteobacteria, as *spf* were exclusively found in this bacterial class. This was done as follows: Representative *spf* sequences from all *spf*-containing genera were used as queries in blast searches. All blast “hits” had a low E-value (i.e., high statistical support; typically below 1e-11). In other words, *spf* was identified with a high degree of confidence, or, *spf* was not found. In one case a hit with a poor E-value was found (0.65). Here, we did a manual inspection to decide the presence/absence of *spf*. First, the NCBI Sequence Viewer (<http://www.ncbi.nlm.nih.gov/projects/sviewer/>) was used to locate the intergenic region between *polA* and *engB* (genes that are known to flank *spf*). Next, a manual text search revealed the presence of highly conserved 5' hairpin, and thereafter the entire *spf*. The presence/absence of *spf* in all complete genomes from gamma-proteobacteria is provided in **Table S1**. The presence of *spf* was next mapped on the tree of life, which was produced using the iTol web tool.²³

Alignments and nucleotide diversity

The sequences from the Rfam list and the newly discovered sequences of *spf* were automatically aligned and manually examined using Jalview.²⁴ An alignment containing only one version of each nucleotide variation of *spf* (no redundant *spf* sequences) was used to examine the variations on nucleotide level between families, genera and species. A consensus *spf* sequence was made based on the alignment and was mapped onto an *E. coli* secondary structure (**Fig. 2**).¹ The *spf* alignment in Rfam includes the first 10 nucleotide upstream of the 5' end of *spf*. However, the promoter region of *spf* was not considered in this work, and was not included in the alignment. Existing literature on experimentally verified mRNA targets of Spot 42 were used to map mRNA targets onto the secondary structure of Spot 42 (**Fig. 3**).^{4, 14, 15}

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Supplemental Material

Supplemental data for this article can be accessed on the publisher's website.

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Figure legends

Figure 1

The natural distribution of *spf*. *spf* is restricted to five orders of gamma-proteobacteria (shown in bold letters), four of which share the same closest common ancestor (indicated by an arrow). The circular phylogenetic tree (made using the iTol web tool) shows all major branches of Bacteria. The gamma-proteobacteria phylogeny in the right panel is based on Gao et al.²⁵ Here, numbers in parentheses indicate the number of complete genomes that contain *spf* (first number) and the total number of available complete genomes (second number) in each order. In addition, *spf* is found in 8 Chromatiales draft genomes (asterisk).

Figure 2

Secondary structure consensus model of the Spot 42 RNA. The structure model was made by aligning all known *spf* sequences, and by mapping the consensus sequence onto a secondary structure model of the *E. coli* Spot 42 (based on Møller et al.¹). The structure consists of a relatively long 5' hairpin, a 9 nt long single-stranded region followed by a second hairpin and a rho-independent terminator. Level of identity is shown using different type of letters in the structure. Uppercase bold letters indicate 80–100 % identity, uppercase regular letters indicate 60–79% identity, and lowercase letters indicate <60% identity. Structural segments with family-specific (i.e., *Vibrionaceae*, *Aeromonadaceae* and *Shewanellaceae*) variations are shown in separate colored boxes. Here, circles indicate U or A insertions (compared to the “consensus”). Grey square around a letter symbolizes aberration from the consensus structure.

Figure 3

Potential base-pairing between the Spot 42 RNA and experimentally verified mRNA targets from the following genes: (A) *galK*, (B) *pirin*, (C) *fucI*, (D) *xylF* and *sthA*, (E) *gltA* and *srlA* and (F) *nanC*, (G) *paaK*, *ascF*, *caiA* and *fucP*, (H) *atoD* and *puuE* and (I) *nanT*. **Fig. 3** is based on data from Møller et al.,¹ Hansen et al.,⁴ Beisel and Storz,¹⁴ and Beisel et al.¹⁵

Figure 4

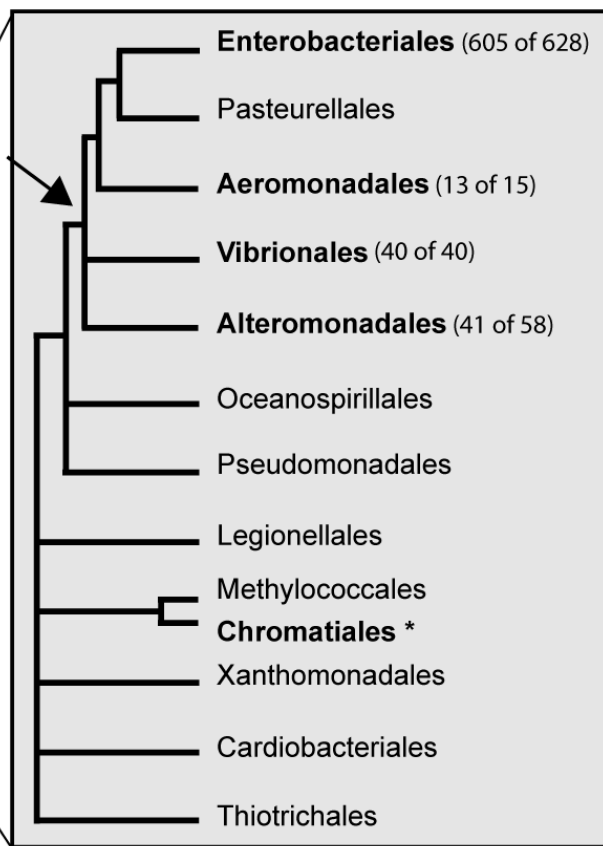
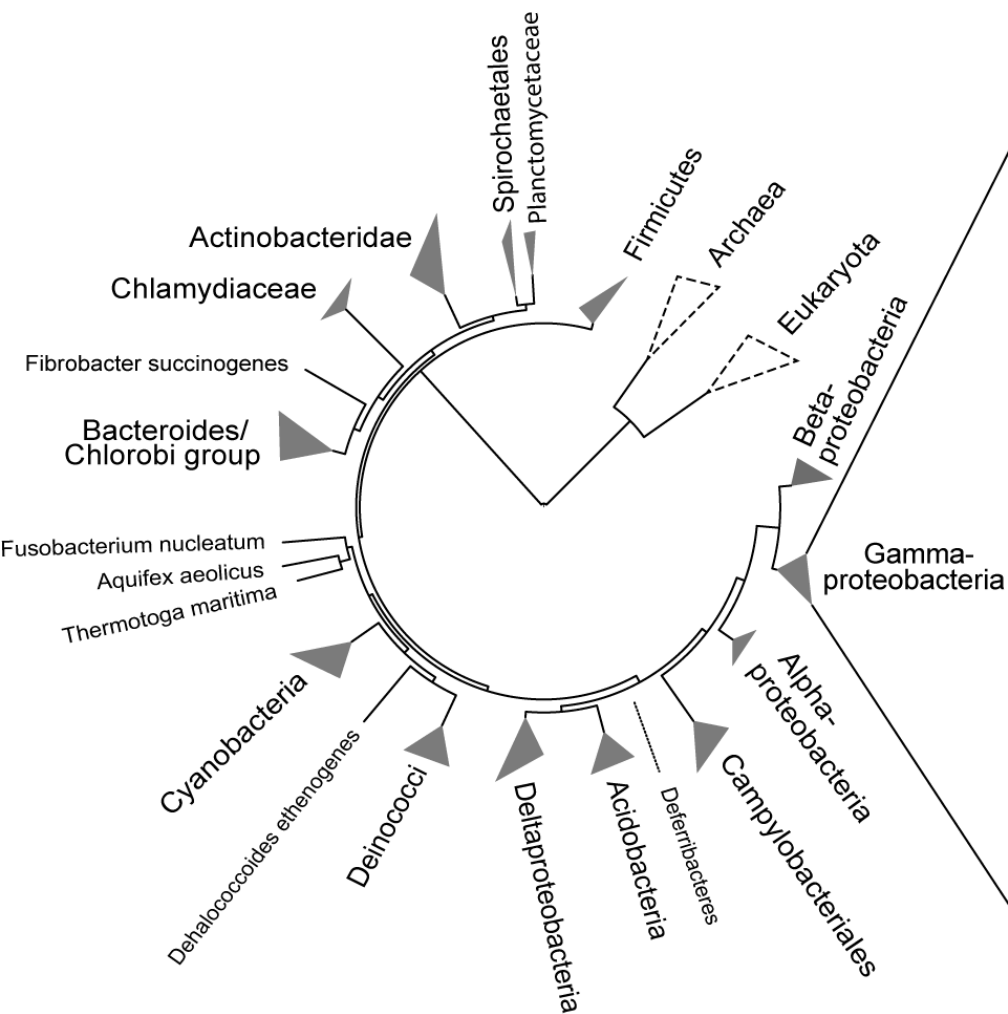
sRNA genes in the intergenic region downstream of *polA*. The figure shows currently known sRNA genes which have been found in the same intergenic region as *spf*. The scale bar shows distance in nucleotides. (A) Representative species containing *spf* are shown. The *VSsrna24* sRNA gene is located downstream of *spf* in *V. cholerae* and *A. salmonicida*. Question mark

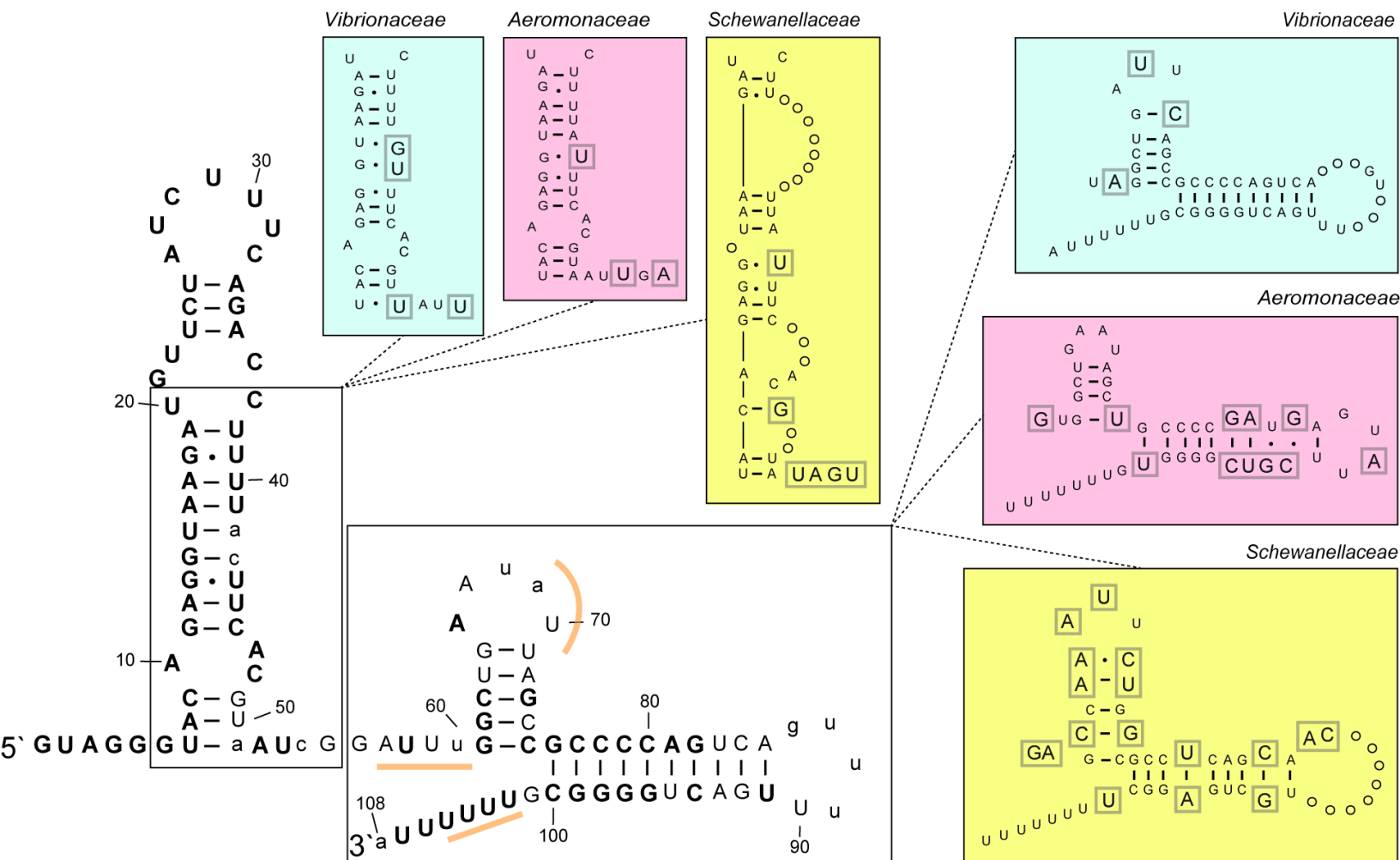
393 denotes hypothetical protein. (B) Genomic location of the sRNA genes *ersA* in *Pseudomonas*
394 *aeruginosa*, *sX13* in *Xanthomonas campestris* and *SMc02857* in *Sinorhizobium meliloti*.

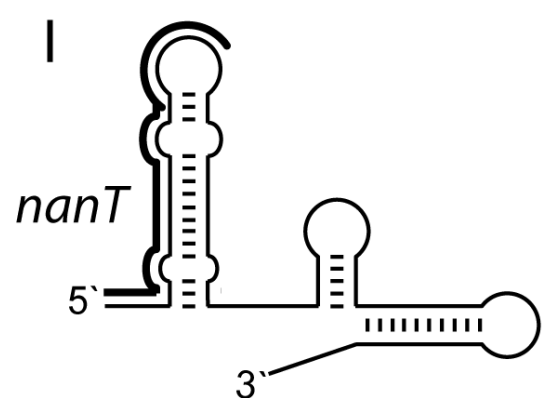
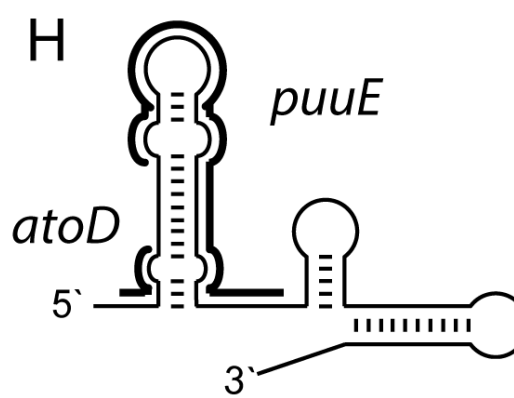
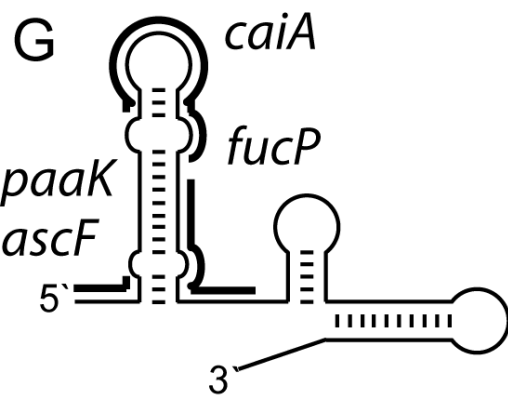
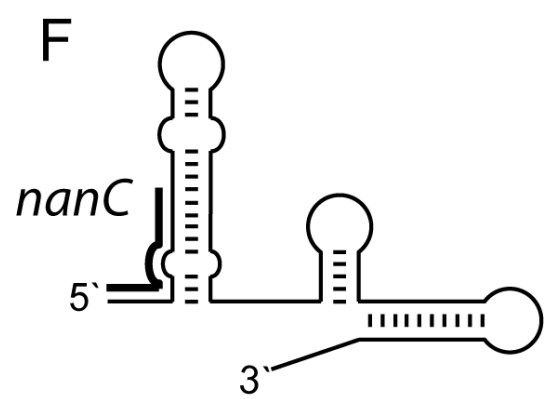
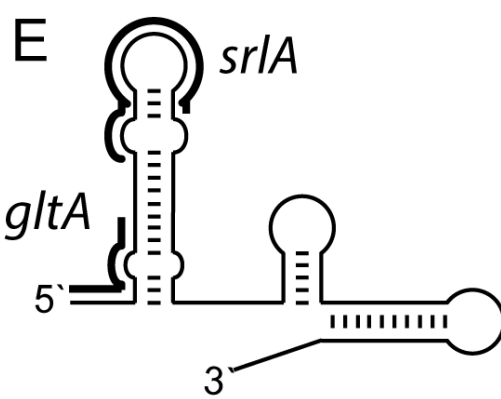
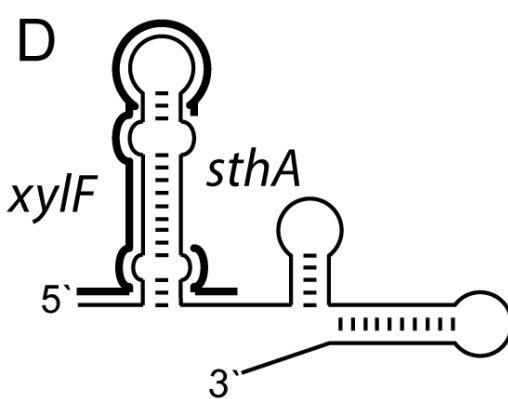
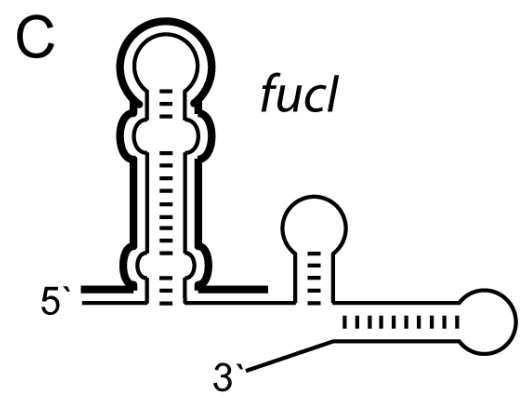
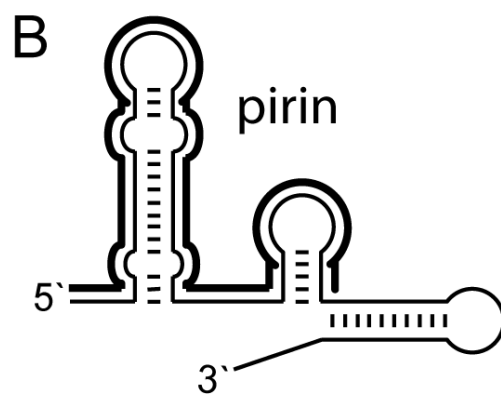
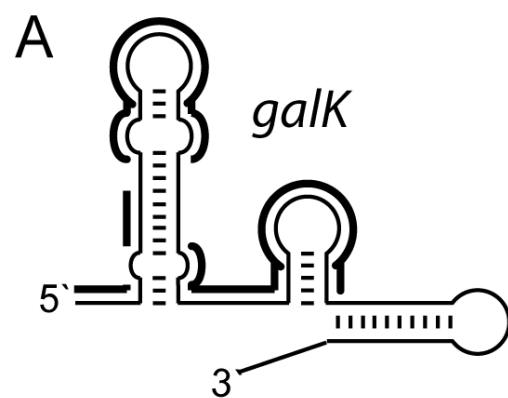
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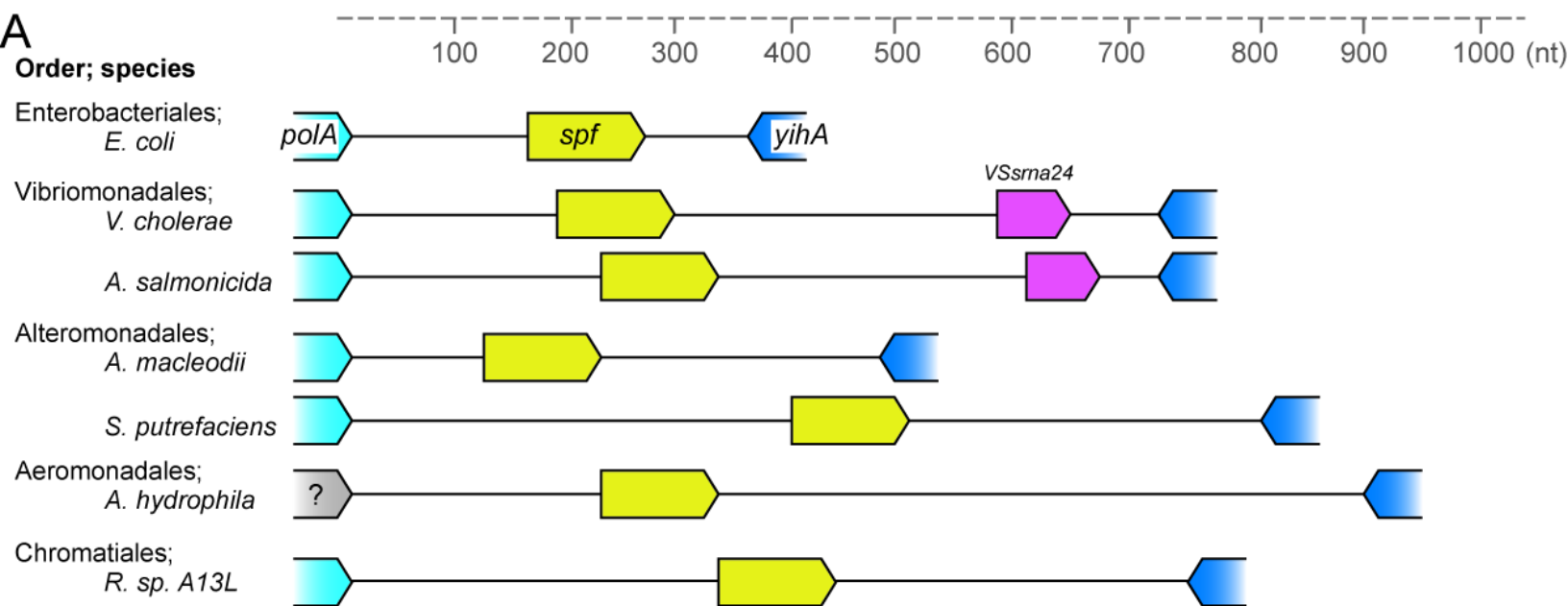






A

Order; species



B

